Bioavailability of Two Oral-Granule Formulations of a Single Dose of 4-Mg Oral Granules of Montelukast: A Randomized, Two-Period Crossover Comparison in Healthy Mexican Adult Volunteers

Gabriel Mendoza-Tamayo¹, Alejandra Rosete-Reyes¹, Jessica González-Bañuelos², Carolina Pérez-Pineda², Ericka López-Bojórquez², Victoria Burke-Fraga¹ and Mario González-de la Parra³

¹CIF-BIOTEC, Médica Sur, S.A.B. de C.V., Mexico City, Mexico City, Mexico
²BioKinetics, S. A. de C. V., Mexico City, Mexico

Abstract

Montelukast is a leukotriene receptor antagonist. In Mexico, the oral granules are indicated for children for the treatment of asthma and allergic rhinitis [3]. The 4 mg oral granules dosage form has been shown to provide systemic exposure similar to that of the 10 mg tablet in adults [4].

Knoor et al. [5] have demonstrated that the montelukast 4 mg oral granules formulation is bioequivalent to the 4 mg chewable tablet formulation when administered in the fasted state. In addition, they evaluated the effect of food on the pharmacokinetics of the oral granules formulation in two studies. In the first study, they found that the administration of a single 4 mg dose of the oral granules formulation with applesauce, a food commonly consumed by young children, had no effect on overall drug exposure, as indicated by the nearly identical AUCₘₙₚ values observed when the oral granules formulation was administered with and without applesauce.

In the second study, they evaluated the effect of the oral granules formulation after consumption of a high-fat breakfast; the findings of this study were that the consumption of a high-fat breakfast had no evident effect on the extent of absorption, as indicated by the similarity between AUCₘₙₚ values after administration of the oral granules formulation with and without food. However, the Cₘₙₚ was statistically significantly lower (36%) and the Tₘₚ delayed (from 2 to 6 hours) after administration with a high-fat breakfast. It is important to point out that the authors acknowledged that it is unlikely that a meal with such a high-fat and calorie content would be consumed by 6 months to 2-year-old patients, and that the delay in absorption should not be regarded as clinically important for a chronically administered drug like montelukast, whose pharmacological effects are more dependent on extent of systemic exposure (i.e., AUC) rather than on maximum peak plasma concentrations [5].

The sponsor of this study (Laboratorios Liomont, S. A. de C. V.) was interested in obtaining the marketing authorization for montelukast 4 mg, as a granule formulation (test formulation) in Mexico (Everest, Laboratorios Liomont, S. A. de C. V., Mexico City, Mexico).

Therefore, the aim of this study was to compare the bioavailability and to determine the bioequivalence of a test formulation containing 4 mg of montelukast, with its corresponding reference drug formulation (Singulair® granules. Merck Sharp & Dohme de Mexico, S. A. de C. V.).

A search of PubMed, MEDLINE and Google data bases for literature published up to May of 2014, using the combination terms montelukast, bioequivalence, bioavailability, pharmacokinetics, oral-granules, Mexico, Mexican and population, did not identify any published data concerning the bioavailability of oral montelukast in the Mexican population.

Keywords: Montelukast; Bioequivalence; Bioavailability; Oral-granules

Introduction

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Materials and Methods

*Corresponding author: Mario González-de la Parra, PvdD. Jesus del Monte No. 77, Col. Cuajimalpa, 05600 Mexico D.F., Mexico, E-mail: mdelparra@biokinetics.com.mx

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The study protocol (P456S026V004) and the informed-consent form were reviewed and approved by an independent ethics and research committee (Comité de Ética e Investigación para Estudios en Humanos, Mexico City, Mexico) on November 26, 2012, and by COFEPRIS (Federal Commission for Protection against Sanitary Risks, for its Spanish acronym) on January 31, 2013. The study was conducted in accordance with the ethical principles of the Declaration of Helsinki and its amendments and the International Conference on Harmonisation for Good Clinical Practice Guideline.

The principal investigator informed the subjects of all procedures, duration of the study, anticipated risks and discomfort it could entail, and an individual written informed consent was obtained prior to the initiation of the study. The study was conducted from April to June of 2013.

Inclusion/exclusion criteria

Healthy Mexican adults aged 18 to 55 years and of either gender were eligible for inclusion in the study. Subjects were recruited from the volunteer’s database at the Center of Pharmacological and Biotechnology Research (clinical unit) in Medica Sur Hospital, Mexico City, Mexico.

Each potential subject had a clinical health evaluation. Classification of subjects as healthy was based on unrewardable findings obtained in a personal interview; a complete physical examination (blood pressure [BP], heart rate, weight, height, temperature and respiratory rate); and diagnostic testing that included a 12-lead ECG, chest radiography, and laboratory blood chemistry tests, serological tests for hepatitis B and C and HIV antibodies, urinalysis, alcohol test, drug-abuse tests and a pregnancy test in women. Other exclusion criteria were alcohol consumption (abstinence during the entire duration of the study), and tobacco consumption (abstinence for at least 48 hours prior to the initiation of the study and during its entire duration).

Systolic and diastolic BP was measured with a sphygmomanometer (Tycos; Welch Allyn, Skaneateles Falls, NY). The BP cuff was applied to the right arm and the reading was taken with the subject in a seated position. Candidates were excluded if laboratory values were significantly out of the reference range and/or if all tests had not been completed. Laboratory testing was performed at Medica Sur Hospital, which has been certified by the Mexican government and the College of American Pathologists. The scope of the certifications included the tests relevant to this study. Before the enrollment of the participants, a personal interview; a complete physical examination (blood pressure, heart rate, weight, height, temperature and respiratory rate); and a pregnancy test in women. Other exclusion criteria were alcohol consumption (abstinence during the entire duration of the study); and tobacco consumption (abstinence for at least 48 hours prior to the initiation of the study and during its entire duration).

Study design and drug administration

A single-dose randomized-sequence, single-blind, two-period crossover design under fasting conditions was used. The subjects were admitted to the clinical site on the day before the drug administration, and were randomly assigned by a pharmacist in the presence of quality assurance personnel at the clinical unit in a 1:1 ratio, using a computer-generated table of random numbers to one of the two sequences (test formulation containing montelukast sodium equivalent to 4 mg of montelukast (lot 239D0017; expiration date January 2015) followed by the reference formulation containing montelukast sodium equivalent to 4 mg of montelukast (lot H013658; expiration date April 2014), or vice versa.

To ensure reliable baseline plasma measurements, participants underwent a 10 hour overnight fast. Based on the reported mean plasma half-life of montelukast, which ranges from 2.7 to 5.5 hours [3], a seven-day washout period was considered appropriate because it exceeds the seven half-lives required by COFEPRIS [6].

Blood samples were drawn for baseline plasma determinations in the following way. An 18-GA x 1.16 in (1.3 x 30 mm) indwelling angiocatheter (BD-InSyte, Becton Dickinson Ind. Cir. Ltda, Minas Gerais, Brazil) was inserted in a suitable forearm vein and a 7.5 mL blood sample was drawn into a heparin-treated vacuum tube (S-Monovette, Sarstedt AG & Co., Nümbrecht, Germany).

Subjects were administered a sachet containing 4 mg oral granules of the test or the reference formulation to be placed in the oral cavity and ingested with the aid of 250 mL of water. Additional blood samples were drawn at 0.25, 0.5, 1, 1.5, 2, 2.5, 3, 3.5, 4, 5, 6, 8, 10, 12 and 24 hours after administration.

During hospitalization, the subjects were under medical surveillance, and during the washout period participants maintained contact with the investigators to report any adverse events (AEs).

Plasma was obtained by centrifugation (3000 rpm for 15 minutes at room temperature) and stored at -70°C ± 10°C (until it was transported to the analytical unit where it was stored at -75°C ± 5°C until it was analyzed). After a seven-day washout period, participants returned to the clinical unit, where the alternative formulation was administered as in the first treatment period.

Subjects were asked to refrain from water and food intake for three hours after the study drug administration. Their diet, for each treatment period, consisted of three standardized meals (2113 kcal/d) at 3.25, 6.5 and 12.75 hours after the study drug administration.

Determination of montelukast plasma concentrations

Chemicals: Montelukast sodium (lot: 10-202, purity 98.2%) secondary standard was obtained from Sintenovo (Estado de Mexico, Mex.) and loratadine reference standard from the USP (Rockville, MD). All solvents (including water) were HPLC-mass spectrometric grade (Avantor Performance Materials, Inc., Phillipsburg, NJ) and all reagents were analytical grade (Mallinckrodt Baker, Inc., Phillipsburg, NJ).

Method and sample preparation: Montelukast sodium plasma levels were determined as montelukast (the free acid form of the montelukast sodium salt) by using a HPLC method coupled with mass spectrometry (MS/MS) developed and validated by personnel of Biokinetics in Mexico City, Mexico. The method included the following: 250 µL of plasma, 10 µL of internal standard (loratadine, 0.05 µg/mL) and 750 µL of acetonitrile. These components were vortexed in a 2.0-mL conical tube (Sarstedt AG & Co.) for one minute. The tube was centrifuged at 8000 rpm for five minutes at room temperature (25°C). The supernatant was separated and injected (volume of injection = 3 µL) into the chromatographic system (HPLC, Agilent Technologies, model 1200, Palo Alto, California).

Chromatographic Conditions: Montelukast concentrations were determined with a 50 × 4.6-mm internal-diameter column of 1.8 µm particle size (Zorbax’ Eclipse XDB-C8, Agilent Technologies) and eluted with a mobile phase consisting of a mixture (30:70 v/v) of an aqueous buffer solution (ammonium formate, 10 mM; pH 6.0 ± 0.1) and acetonitrile. The column temperature was 25°C. Flow rate was maintained at 0.6 mL/minute and the montelukast and loratadine were detected by a triple-quadrupole mass spectrometer (Agilent Technologies, model G6410B). The spectrometric (MS/MS) analysis was performed by monitoring the transitions 586.2→422.3 m/z for montelukast and 383.3→337.2 m/z for loratadine (IS). The
spects of the study, concerning the occurrence AEs. Subjects were asked to spontaneously report any AE to the investigators at any time during the study, including the washout period. Data for all AEs were recorded on a case-report form.

AEs that were life-threatening, led to death, hospitalization, disability, and/or medical intervention to prevent permanent impairment or damage, were considered serious.

Pharmacokinetic and statistical analyses: Sample size calculation [8] was based on the within-subject variability of montelukast $C_{\text{max}}$ with a % CV of 21.2% [5]. This calculation was performed considering the following values: 1 - $\beta = 0.8$, $\alpha = 0.05$, % CV = 21.2, and an equilibrium range of 80% to 125%, yielded with a sample size of 22 subjects. In this study, a sample size of 26 subjects was used because, at the time the study was conducted, a minimum sample size of 24 subjects was required for bioequivalence studies by COFEPRIS.

Individual plasma concentration–time curves were constructed; $C_{\text{max}}$ and $T_{\text{max}}$ were directly obtained from these curves, the area under the plasma concentration-time curve from time baseline to the last measurable concentration ($AUC_{0-t}$) was calculated by a non-compartmental method using the trapezoidal rule. From the terminal log-decay phase, the elimination rate constant ($k_e$) was estimated using linear regression, and the $t_{1/2}$ was estimated using the following equation [8]:

$$t_{1/2} = \ln 2 / k_e,$$

where $L$ was defined as the natural logarithm. Extrapolation of AUC from baseline to infinity ($AUC_{0-\infty}$) was calculated as follows:

$$AUC_{0-\infty} = AUC_{0-t} + Ct/ke,$$

where $Ct$ was the last measurable plasma concentration.

To assess the bioequivalence between the test and reference formulations, $C_{\text{max}}$, $AUC_{0-t}$, and $AUC_{0-\infty}$ were calculated using log-transformed data for these parameters, was carried out at the 5% significance level ($\alpha = 0.05$).

The 90% CIs of the geometric mean ratios (test/reference) of $C_{\text{max}}$, $AUC_{0-t}$, and $AUC_{0-\infty}$ were calculated using log-transformed data. The test and the reference formulations were considered bioequivalent if the 90% CIs of these parameters fell within a predetermined range of 80% to 125%. All pharmacokinetic and statistical analyses were performed using WinNonlin Version 5 (Pharsight, Mountain View, California).

Results

Table 1 show the demographic characteristics for a total of 26 subjects, who were enrolled and completed the clinical stage of the study.

Pharmacokinetic parameters

Mean plasma concentration-time curves of the two formulations are shown in Figure 1. This figure suggests comparable mean plasma concentration-time curves.

Table 2 shows the pharmacokinetic parameters ($C_{\text{max}}$, $T_{\text{max}}$, $t_{1/2}$, $AUC_{0-t}$, and $AUC_{0-\infty}$) for both formulations.

No significant period or sequence effects were detected based on the ANOVA of $C_{\text{max}}$, $AUC_{0-t}$, and $AUC_{0-\infty}$ (data not provided).

Table 3 shows the bioequivalence statistics using the log-transformed data of $C_{\text{max}}$, $AUC_{0-t}$, and $AUC_{0-\infty}$: geometric mean ratios (test/reference), 90% CI, and the intra-subject %CV.

The 90% CIs for montelukast $C_{\text{max}}$, $AUC_{0-t}$, and $AUC_{0-\infty}$ were 97.57% to 109.35%, 101.81% to 108.92%, and 101.55% to 109.96%, respectively. The 90% CIs of the geometric mean ratios of the three parameters fell within the predetermined range of 80% to 125%.

These results indicate that the bioequivalence criteria were met.

Tolerability

Four of the 26 subjects reported a total of 5 AEs. All of them were reported after the administration of the reference formulation: two

In this study of healthy, fasting, Mexican adult subjects, who received a single dose of either the test or reference formulation, it was concluded that the test formulation of 4-mg oral granules of montelukast met the Mexican regulatory requirements to assume bioequivalence, based on the rate and extent of absorption. Both formulations were well tolerated.

Conclusions

In addition, the effect of food was not evaluated because it has been reported that the extent of the absorption of montelukast (as indicated by AUC values) is not affected by food intake, especially when considering the typical diet of young children [5].

Further studies are needed to compare the test formulation with the reference formulation in Mexican patient groups. The results of this study might serve as a reference for future controlled studies of montelukast in a Hispanic population.

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